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Glucose isomérases avec un optimum de pH changé

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#### Description

#### **TECHNICAL FIELD**

- [0001] The present invention relates to the application of protein engineering technology to improve the properties of metalloenzymes. A method for selecting amino acids which upon alteration will influence the pH-activity profile of metalloenzymes is provided. Said method is applied to glucose isomerase. The present invention also provides mutated glucose isomerase molecules with an altered pH optimum. Specifically the acidic flank of the pH-activity profile is shifted towards lower pH.
- [0002] The present invention further provides recombinant glucose isomerases that advantageously can be applied in the production of fructose syrups, in particular high fructose corn syrups.

### **BACKGROUND OF THE INVENTION**

15 Industrial application of glucose isomerase

[0003] In industrial starch degradation enzymes play an important role. The enzyme  $\alpha$ -amylase is used for liquefaction of starch into dextrins with a polymerization degree of about 7-10. Subsequently the enzyme  $\alpha$ -amyloglucosidase is used for saccharification which results in a syrup containing 92-96% glucose. The reversible isomerization of glucose into fructose is catalyzed by the enzyme glucose (or xylose) isomerase. The correct nomenclature of this enzyme is D-xylose-ketol-isomerase (EC 5.3.1.5) due to the enzyme's preference for xylose. However, because of the enzyme's major application in the conversion of glucose to fructose it is commonly called glucose isomerase. The equilibrium constant for this isomerization is close to unity so under optimal process conditions about 50% of the glucose is converted. The equilibrium mixture of glucose and fructose is known as high fructose syrup.

25 [0004] Fructose is much sweeter to the human taste than glucose or sucrose which makes it an economically competitive sugar substitute.

[0005] Many microorganisms which were found to produce glucose isomerase, have been applied industrially. A detailed review of the industrial use of glucose isomerases has been given by Wen-Pin Chen in Process Biochemistry, 15 June/July (1980) 30-41 and August/September (1980) 36-41.

30 [0006] The Wen-Pin Chen reference describes culture conditions for the microorganisms, as well as recovery and purification methods for the enzyme. In addition it also summarizes the properties of glucose isomerases such as the substrate specificity, temperature optima and pH optima, heat stability and metal ion requirement.

[0007] Glucose isomerase requires a bivalent cation such as Mg<sup>2+</sup>,Co<sup>2+</sup>, Mn<sup>2+</sup> or a combination of these cations for its catalytic activity. Determination of 3D structures of different glucose isomerases has revealed the presence of two metal ions in the monomeric unit (Farber et al., Protein Eng. <u>1</u> (1987) 459-466; Rey et al., Proteins 4 (1987) 165-172; Henrick et al., Protein Eng. 1 (1987) 467-475).

[0008] Apart from a role in the catalytic mechanism, bivalent cations are also reported to increase the thermostability of some glucose isomerases (M. Callens et al. in Enzyme Microb. Technol. 1988 (10), 695-700). Furthermore, the catalytic activity of glucose isomerase is severely inhibited by Ag+, Hg²+, Cu²+, Zn²+ and Ca²+.

40 [0009] Glucose isomerases usually have their pH optimum between 7.0 and 9.0. There are several reasons why it would be beneficial to use glucose isomerase at a lower pH value. Three of these reasons;

- a) stability of the sugar molecules,
- b) adaptation both to previous and/or later process steps and
- 45 c) stability of the enzyme,

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will be further described below to illustrate this.

- a) Under alkaline conditions and at elevated temperatures the formation of coloured by-products and the production of a non-metabolizable sugar (D-Psicose) are a problem. The desired working pH should be around 6.0. Around this pH degradation of glucose and fructose would be minimal.
  - b) A lowered pH optimum is also desirable for glucose isomerase when this enzyme is to be used in combination with other enzymes, or between other enzymatic steps, for example in the manufacturing of high fructose syrups. In this process one of the other enzymatic steps, the saccharification by  $\alpha$ -glucoamylase is performed at pH 4.5.
  - c) Most of the known glucose isomerases are applied at pH 7.5. This pH value is a compromise between a higher initial activity at higher pH and a better stability of the immobilized enzyme at lower pH, resulting in an optimal productivity at the pH chosen (R. v. Tilburg, Thesis: "Engineering aspects of Biocatalysts in Industrial Starch Conversion Technology", Delftse Universitaire Pers, 1983). Application of glucose isomerase at a pH lower than 7.5

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could benefit from the longer half-life and, combined with an improved higher specific activity, would consequently increase the productivity of the immobilized enzyme at that lower pH.

From the above it can be concluded that there is need for glucose isomerases with a higher activity at lower pH values under process conditions.

[0010] Many microorganisms were screened for a glucose isomerase with a lower pH optimum. Despite many efforts, this approach did not lead to novel commercial glucose isomerases.

[0011] In order to be able to change pH-activity profile of glucose isomerases towards lower pH by protein engineering it is important to recognize the underlying effects which give rise to the rapid decrease in catalytic performance at acidic pH.

#### The role of metal ions in enzymes

[0012] Two different functions for metal ions in enzymes can be envisaged.

15 [0013] First of all metal ions can have a structural role. This means that they are involved in maintaining the proper 3D-structure and, therefore, contribute to the (thermo)stability of the enzyme molecule. An example of such a structural and stabilizing role is Ca<sup>2+</sup> in the subtilisin family of serine proteinases.

[0014] Secondly, metal ions can act as a cofactor in the catalytic mechanism. In this case the enzyme activity is strictly dependent upon the presence of the metal ion in the active site. The metal ion may for instance serve as a bridge between the enzyme and the substrate (e.g. Ca<sup>2+</sup> in phospholipase binds the phosphate group of the substrate) or it may activate water to become a powerful nucleophilic hydroxyl ion (Zn<sup>2+</sup>-OH<sup>-</sup>).

[0015] Examples are the Zn<sup>2+</sup>-proteases such as thermolysin and carboxypeptidase, carbonic anhydrase (Zn<sup>2+</sup>), phospholipase-A<sub>2</sub> (Ca<sup>2+</sup>) staphylococcal nuclease (Ca<sup>2+</sup>) and alkaline phosphatases (Mg<sup>2+</sup>, Ca<sup>2+</sup>). Examples of alpha/ beta barrel enzymes which require cations to polarize a carboxyl or a carbonyl group in order to transfer hydrogen are glucose/xylose isomerase (Mg<sup>2+</sup>), ribulose-1,5-biphosphate carboxylase/oxygenase (RU-BISCO) (Mg<sup>2+</sup>), enolase (Mg<sup>2+</sup>), yeast aldolase (Mg<sup>2+</sup>, K<sup>1+</sup>), mandolate racemase (Mg<sup>2+</sup>), muconate cycloisomerase (Mn<sup>2+</sup>). In the presence of metal chelating agents (such as EDTA), these enzymes loose their activity completely.

[0016] The binding of metal ions in a protein molecule usually involves coordination by 4 or 6 ligands. Depending on the type of metal ion, different ligands are found. For instance magnesium and calcium are usually liganded by oxygen atoms from either a carbonyl group of the protein main chain, a carbonyl group from a glutamine or asparagine side chain or the carboxylate from an aspartic- or glutamic acid side chain. Zinc and copper ions are usually liganded by nitrogen atoms from a histidine side chain or the sulfur atoms from cystein and methionine.

#### Factors determining the pH dependence of an enzyme

[0017] The activity of an enzyme is dependent on the pH value of the aquaous medium. This dependence is caused by the (de)protonation of ionizable groups in the active site of the enzyme on the one hand, and ionizable groups of the substrate, or product (if present) on the other hand. Ionizable groups in proteins involve the side chains of the basic amino acids lysine, arginine and histidine (carrying a positive charge in the protonated form), and the acidic amino acids aspartic acid, glutamic acid, cystein and tyrosine (all carrying a negative charge upon deprotonation). Furthermore, the amino group of the N-terminus and carboxyl group of the C-terminus carry a positive and negative charge respectively. The pK<sub>a</sub>-values of some amino acids are depicted in Table 1.

Table 1.

|                       |            | рК <sub>а</sub> |
|-----------------------|------------|-----------------|
| Positive charge (base |            |                 |
|                       | N-terminus | 7.5-8.5         |
|                       | Lysine     | 10.5            |
|                       | Arginine   | 12.5            |
|                       | Histidine  | 6.0-7.0         |
| Negative charge (acid | )          |                 |
| <del></del>           | C-terminus | 3.0-4.0         |

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Table 1. (continued)

| Ionizable groups of amino acids as occurring in proteins [Cantor and Schimmel, 1980, Biophysical chemistry, W.H. Freeman, San Fransisco] |                 |  |  |  |
|--|-----------------|--|--|--|
|  | рК <sub>а</sub> |  |  |  |
| Negative charge (acid)   |                 |  |  |  |
| Aspartic acid  | 3.9             |  |  |  |
| Glutamic acid  | 4.3             |  |  |  |
| Cystein  | 8.3             |  |  |  |
| Tyrosine   | 10.1            |  |  |  |

[0018] It should be realised that these  $pK_a$ -values are valid for model compounds and that great variations both within and between different proteins occur, due to the specific environment of the ionizable group. Electrostatic effects are known to play a fundamental role in enzyme function and structures (see J.A. Matthew et al, CRC Critical Reviews in Biochemistry, 18 (1985) 91-197). The presence of a positive charge near an ionizable group will lower its  $pK_a$  while a negative charge will cause an increase in  $pK_a$ . The magnitude of the effect decreases with the distance between the ionizable group and the charge. Moreover, the magnitude of this decrease is dependent upon the dielectric constant of the medium. Especially catalytic residues may reveal  $pK_a$ -values which deviate from these averages (see for instance Fersht, Enzyme structure and mechanism, 1985, W.H. Freeman, New York).

[0019] The pH-dependence of an enzyme catalyzed reaction can be dissected into the pH-dependence of the Michaelis constant  $K_{m}$  and the pH-dependence of the turn-over rate constant  $K_{cat}$  (equivalent to  $V_{max}$ ). These parameters represent the binding of the substrate in the ground-state and transition-state respectively. The pH-dependent (de) protonation of amino acid side chains which affect the binding of both substrate forms, or which are otherwise involved in the catalytic event (e.g. proton uptake and release as in general base catalysis), therefore, determine the pH-activity profile of an enzyme.

[0020] For instance the protonation of the histidine in the catalytic triad of serine proteases (both the trypsin- and subtilisin family) is responsible for the loss of activity at lower pH-values (<7). In this case, the  $pK_a$  of the enzyme activity is directly related to the  $pK_a$  of this histidine residue.

[0021] As a second example, the two aspartic acid residues in aspartyl proteases, such as pepsin and chymosin, can be mentioned. These groups determine the pH optimum of these proteases. The typical structural arrangement of the aspartic acids causes them to have different pK<sub>a</sub>-values leading to the bell-shaped pH-activity profile.

[0022] It is known that altering the surface charge by extensive chemical modification can lead to significant changes in the pH dependence of catalysis. However in many cases this approach leads to inactivation and/or unwanted structural changes of the enzyme because these methods are rather unspecific. Selective chemical modification of lysines in cytochrome c was shown to have an effect on the redox-potential (D.C. Rees, J. Mol. Biol. 173, 323-326 (1980)). However, these results have been criticized because the bulky chemical reagent used for modification could perturb the structure of the protein.

[0023] Using the 3D-structure of a protein to anticipate the possibility of structural perturbation and site-directed mutagenesis, it is possible to modify the charge distribution in a protein in a very selective way.

[0024] Fersht and coworkers have shown that it is possible to manipulate the pH-activity profile of subtilisin by site-directed mutagenesis (Thomas et al, Nature,  $\underline{318}$ , 375-376 (1985); Russell et al, J. Mol. Biol.,  $\underline{193}$ , 803-813 (1987); Russel and Fersht, Nature  $\underline{328}$ , 496-500 (1987)). Introduction of negatively charged groups at 10-15Å from the active site at the protein surface raises the pK<sub>a</sub> value of the active site histidine. Conversely, making the surface more positively charged lowers the pK<sub>a</sub> of the acidic groups. Changing either Asp99 at 13 Angstroms or Glu156 at 15 Angstroms from the active site to a lysine lowers the pK<sub>a</sub> of the active site histidine by 0.6 pH units. Changing both residues simultaneously to give a double mutant with a change of four charge units, lowers the pK<sub>a</sub> by 1.0 pH unit. It appears that changes in Coulombic interactions can be cumulative.

### Glucose isomerase mutants

[0025] WO 89/01520 (Cetus) lists a number of muteins of the xylose isomerase which may be obtained from <a href="Strep-tomyces rubiglnosus">Strep-tomyces rubiglnosus</a> and that may have an increased stability. The selection of possible sites that may be mutated is based on criteria differing from the ones used in the present invention. More than 300 mutants are listed but no data are presented concerning the characteristics and the alterations therein of the mutant enzyme molecules.

### Methodologies for obtaining enzymes with improved properties

[0026] Enzymes with improved properties can be developed or found in several ways, for example by classical screening methods, by chemical modification of existing proteins, or by using modern genetic and protein engineering techniques.

[0027] Site-directed mutagenesis (SDM) is the most specific way of obtaining modified enzymes, enabling specific substitution of one or more amino acids by any other desired amino acid.

#### Summary of the Invention.

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[0028] The present invention provides methods relating to the production of new mutant metalloenzymes by expression of genes encoding said enzymes having amino acid sequences which differ in at least one amino acid from the corresponding wildtype metalloenzymes and which exhibit altered catalytic properties. Specifically, the pH-activity profile is altered by changing the overall charge distribution around the active site.

[0029] According the invention provides a method for altering the pH-activity profile of a Wild-type glucose isomerase enzyme such that said profile, or the acidic part thereof, is shifted towards a lower pH, comprising the substitution of a negatively charged amino acid residue by a neutral one or by a positively charged one, or the substitution of a neutral residue by a positively charged one, wherein said substitution is at a position selected from the group consisting of: Ala25, Phe61, Gln204, and Leu258; wherein said residues of the glucose isomerase are from Actinoplanes missouriensis whose sequence is shown in Figure 2, or the corresponding position in a homologous glucose isomerase.

[0030] The invention further provides a process for obtaining a mutant glucose isomerase molecule which comprises: (a) obtaining a DNA sequence encoding a glucose isomerase; (b) mutating this sequence at selected nucleotide positions to provide an enzyme altered at any one of positions Ala25, Phe61, Gln204, and Leu258 by a substitution of a negatively charged amino acid residue by a neutral one or by a positively charged one, or the substitution of a neutral residue by a positively charged one, wherein said residues of the glucose isomerase are from Actinoplanes missouriensis whose sequence is shown in Figure 2, or the corresponding position in a homologous glucose isomerase; (c) cloning the altered sequence into an expression vector in such a way that the DNA sequence can be expressed; (d) transforming a host organism or cell with said vector; (e) culturing said host organism; and, (f) isolating the altered glucose isomerase.

30 [0031] Such a process may further comprise the use of the mutant glucose isomerase in the production of a fructose syrup.

[0032] In the above aspects of the invention, the wild-type glucose isomerase may be from a microorganism of the order Actinomycetales, such as Actinoplanes missouriensis, and the amino acid replacements may be in particular A25K, F61K, Q204K, and L258K.

35 [0033] When the glucose isomerase is the <u>Actinoplanes missouriensis</u> isomerase of Figure 2 the replacement may be F61 K+K253R.

### Brief Description of the Figures.

#### 40 [0034]

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Figure 1 shows a schematic representation of the active site of glucose isomerase from Actinoplanes missouriensis derived from the three dimensional structure of the glucose isomerase - xylitol complex. The inhibitor is shown in full detail in the centre of the figure. For the amino acid residues only those atoms are drawn which are involved in hydrogen bonding. Amino acid residue names are in boxes drawn with solid lines, solvent molecules are in boxes drawn with dashed lines. Metal binding sites are indicated by ovals numbered 395 and 580. Dashed lines indicate electrostatic interactions: the thin dotted lines represent hydrogen bonds, the fat dashed lines the proposed ligation of the metals. Strictly conserved residues are marked by an asterix. For non-conserved residues the substitutions found are indicated.

Figure 2 shows the alignment of amino acid sequences of glucose isomerases from different sources. The complete sequence of <u>Actinoplanes missouriensis</u> glucose isomerase is given. The amino acid sequence of <u>Ampullariella</u> glucose isomerase differs from that of the published sequences (Saari, J. Bacteriol., 169, (1987) 612) by one residue: Proline 177 in the published sequence was found to be Arginine.

The <u>Streptomyces thermovulgaris</u> sequence has only been established up to amino acid 346. Undetermined residues are left blank. A dot indicates the absence of an amino acid residue at this position as compared to any of the other sequences. The different species are indicated by the following symbols:

Ami.: Actinoplanes missouriensis DSM 4643
Amp.: Ampullarella species ATCC31351
Svi.: Streptomyces violaceoruber LMG 7183

Smu.: Streptomyces murinus

Sth.: Streptomyces thermovulgaris DSM 40444

Art.: Arthrobacter species
Bsu.: Bacillus subtilus
Eco.: Escherichia coli
Lxy.: Lactobacillus xylosus

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The secondary structure assignment was made in the structure of <u>Actinoplanus missouriensis</u>. The helices in the barrel are enclosed by solid lines. The  $\beta$ -strands are in the shaded boxes.

Figure 3 shows the pH-activity profile of glucose isomerase in the presence of 200 mM xylose and 10 mM magnesium (squares) and 1 mM manganese (circles).

Figure 4 shows the reaction scheme for the isomerisation catalyzed by glucose isomerase in the presence of metal ions.

E = enzyme, S = substrate, M = metal ion, P = product.

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Figure 5 shows the pH dependence of the reaction of glucose isomerase with xylose and  $Mg^{2+}$  as observed with steady-state experiments.  $K_1$ ,  $K_2$ ,  $K_3$  and  $K_4$  are equilibrium constants explained in the text and in the reaction scheme given in Figure 4.

Figure 6 shows the pH-activity profile for the mutants K294R and K294Q in the presence of 200 mM xylose and 10 mM magnesium.

Figure 7 shows the pH-activity profiles for E186Q and E186D in the presence of 200 mM xylose and 10 mM Mg<sup>2+</sup>.

Figure 8 shows the pH-activity profiles for E186D and E186Q in the presence of 200 mM xylose and 1 mM Mn<sup>2+</sup>.

Figure 9 shows the pH-activity profile of the mutant D255N in the presence of 200 mM xylose and 1 mM manganese.

Figures 10-20 show the normalized pH-activity profiles for the following mutants:

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F254K (Fig.10), F94R (Fig.11), F61K (Fig.12), A25K (Fig.13), D57N (Fig.14), L258K (Fig.15), Q204K (Fig.16), R23Q (Fig.17), H54N (Fig.18), H290N (Fig.19), T95D (Fig.20).

Conditions are mentioned in the Figures.

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Figure 21 shows the normalized pH-activity profile for mutant F61KK253R. Conditions are mentioned in the Figure.

### **DETAILED DESCRIPTION OF THE INVENTION**

- [0035] The present invention describes the modification of enzymes to improve their industrial applicability. The invention makes use of recombinant DNA techniques. Such techniques provide a strong tool for obtaining desired amino acid replacements in the protein of choice. Because of the virtually unlimited amount of possible amino acid replacements it is preferable to use a selective approach. The present approach relies on the well coordinated application of protein crystallography, molecular modelling and computational methods, enzymology and kinetics, molecular biology and protein chemistry techniques. The strategies for the identification of targeted mutations are innovative in the sense that it is recognized that point mutations rarely cause only local perturbations. Mutations generally affect several different properties of the protein at once. Therefore, although the disclosed strategies make use of well established structure-function relationships, they also provide a rational way to avoid or correct unwanted alterations of secondary properties.
- 55 [0036] Extensive biochemical investigation of the designed mutants results in the identification of mutants with improved properties.
  - [0037] By 'improved properties' as used herein in connection with the present glucose isomerase enzymes we mean enzymes in which the acidic flank of the pH-activity profile shifted towards a more acidic pH optimum relative to the

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corresponding wildtype enzymes.

[0038] It was established that the pH-activity profile of wildtype glucose isomerase reveals a decrease in activity at both acidic pH below 7.0 and at alkaline pH beyond pH 8.0 (Example 1 - Figure 3). As discussed earlier it would be preferable to use glucose isomerase at lower pH.

[0039] Surprisingly, it was found that the drop in activity at the acidic side of the pH-activity profile is caused by the protonation of one or more amino acid side chains, which are directly involved in the coordination of the catalytic metal ion of glucose isomerase. This was deduced from the fact that the apparent association constants for metal binding, as determined by steady-state kinetics, showed a similar pH-dependency (Example 1 - Figure 5). This can be described by the following model:

site + Mg<sup>2+</sup> K<sub>sss</sub> site:Mg<sup>2+</sup>

pK<sub>s</sub> 1|
site:H<sup>2</sup>

in which "site" refers to the geometrical binding site composed of the different (ionizable) ligands. The enzyme is only active when the site is occupied by an Mg<sup>2+</sup>-ion, which, in turn, can only bind to the unprotonated metal binding site ("site") and not the protonated one ("site:H+"). The pH-dependent protonation of the metal binding site is characterized by the pK<sub>a</sub>, whereas the metal binding to the unprotonated site is characterized by the association constant K<sub>ass</sub>. The apparent association constant for metal binding is a function of the true K<sub>ass</sub>, the pK<sub>a</sub> and the pH and can be described as follows:

 $K_{ass}^{apparent} = K_{ass} * (1 + 10^{pKa-pH)})^{-1}$ 

[0040] The rate of the reaction is proportional to the fraction of the enzyme molecules which is complexed with  $Mg^{2+}$ . This fraction increases at a higher  $[Mg^{2+}]^*K_{ass}^{apparent}$ .

[0041] Although the presented model was described after an observation made in glucose isomerase, it is obviuous that this model, and the method derived therefrom, can be used for metalloenzymes in general, provided that the inactivation at low pH is due to the dissociation of the metal ions.

[0042] The drop in activity observed at alkaline pH is due to a decrease in the maximal velocity ( $V_{max}$ ), reflecting the deprotonation of an amino acid residue that is essential for the catalytic mechanism.

[0043] From the model described above, which can be used to explain the decrease in activity at the acidic side of the pH-activity profile, it can be deduced that in order to increase the activity of glucose isomerase at lower pH values,  $K_{ass}$  has to be increased and/or the p $K_a$  has to be decreased.

[0044] Therefore, DNA sequences coding for metalloenzymes such as glucose isomerases may be mutated in such a way that the mutant proteins reveal a change in the pH-activity profile as a result of a change in  $pK_a$  of amino acid side chains acting as ligands in the metal binding.

Shift of the pH-activity profile of metalloenzymes to lower pH, by changing the pK<sub>a</sub> of amino acid side chains

- 45 [0045] In order to shift the pK<sub>a</sub>'s of metal coordinating ligands to a more acidic pH, residues have to be introduced which increase the overall positive charge around the metal binding site of metalloenzymes. Consequently, the pH dependence of the activity of metalloenzymes, for which the activity at the acidic side of the pH optimum is caused by the pK<sub>a</sub> of metal binding, will change accordingly. This charge alteration will stabilize the negative charge of the ionizable groups which are responsible for the pH dependence of the metal binding through long range electrostatic effects.
  - [0046] According to a preferred embodiment the shift of the pH-activity profile of glucose isomerase to a more acidic pH is achieved by increasing the overall positive charge around the active site of glucose isomerase.

[0047] Around neutral pH a net increase in positive charge can be obtained by:

- replacing a negatively charged residue (Asp or Glu) by a neutral one
- replacing a neutral residue by a positively charged one (Arg or Lys)
- replacing a negatively charged residue by a positively one.

[0048] For the selection of residues, which are suitable to be mutated, the following criteria can be formulated:

I. Select those positions at which substitution will lead to a net increase in positive charge within a 15 Ångstroms radius around the target residues. The target residues are the ionizable groups which are involved in the coordination of the cation.

Eliminate from this collection:

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- All positions that already contain a positive charge: arginine or lysine.
- All positions that cannot harbour an arginine or a lysine because these residues would lead to inadmissible
   Van der Waals overlap with the backbone atoms of the protein.
- All positions at which an arginine or a lysine would need extensive adaptation of additional positions in the direct environment in order to avoid Van der Waals overlap.
- All positions at which substitution into arginine or lysine would lead to a buried uncompensated charge in a hydrofobic cluster.
- All positions at which residues should not be replaced because they are involved in typical structural arrangements such as: salt bridges, packing of helices, stabilization of helices by keeping a negative charge at the start of a helix, initiation of helices, e.g. prolines at the start of a helix, Phi-psi angles which are outside the allowed region for the residue that is going to be inserted.

II. In a preferred embodiment the following amino acids are also eliminated from the collection:

- All residues that are implied in catalysis, cofactor binding (such a metal ions and nucleotides)

- All positions that appear to be strictly conserved among homologous enzymes (if available).

III. Subsequently a priority can be attributed to each possible mutation site. This is done by inspection of the structural environment of the residue, the distance to the 'target' residues, the hydrogen bonding pattern in which the residue at said site is involved and the solvent accessibility.

In order to avoid masking of the electrostatic interactions by counter-ions, introduction of charges at sites which can not be shielded from the target residues by solvent, are to prefer. So in general, due to the difference in dielectric properties between the protein and the solvent, charges, which cannot be solvated completely due to the fact that they are buried in the interior of the protein or partially buried in clefts on the protein surface, are more likely to cause effects than charges that are completely solvated. Moreover, in the case of glucose isomerase, the conversion of glucose into fructose is performed at low ionic strength, and therefore, shielding by counter-ions is expected to interfere less seriously with the newly designed charge-charge interaction in the novel glucose isomerase described within the embodiment of the invention.

Criteria for the assignment of low priority to the above selected sites, when replacing into a positive charge are:

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- The introduction of a positive charge and/or elimination of a negative charge will affect the integrity of the quaternary structure.
- The site is completely solvent accessible so that an introduced charge is expected to be shielded from the target residues by the solvent, which therefore will diminish the effect on pKa of the target residues.

[0049] Likewise, increasing the overall negative charge around the metal binding site will shift the  $pK_a$ 's to more basic pH-values.

Shift of the pH-activity profile of metalloenzymes to lower pH, by increasing Kass for the metal binding

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[0050] In another preferred embodiment of the subject invention the shift of the pH-activity profile to a lower pH is achieved by increasing the  $K_{\rm ass}$ .

[0051] The shift of the pH-activity profile of metalloenzymes to a more acidic pH can be achieved by increasing the association constant for metal binding. The association constant for metal binding can be increased by optimization of the coordination of the metals by the ligands. This may be realised by the introduction of better ligands or by introducing more ligands. Electrostatic interactions can contribute to the association constant for metal binding over much longer distances.

[0052] In another preferred embodiment the acidic flank of the pH-activity profile of glucose isomerase is shifted to lower pH by increasing the association constant for metal binding.

[0053] Glucose isomerase binds two magnesium ions per subunit, resulting in the binding of eight cations per tetramer, thereby increasing the total charge by +16 (see Figure 1). Both binding sites are located at the C-terminal end of the  $\beta$ -barrel. The 'Down' binding sites is located rather deep in the barrel and, in the xylitol complex, directly binds two oxygens from the inhibitor. The second binding site ('Up') is located near the end of the  $\beta$ -barrel, close to the active

site cleft and the subunit interface.

[0054] In general, when a positively charged ion binds to a second particle, the association and dissociation rate constants as well as the overall equilibrium affinity constant will depend upon the charge of the second particle. Repulsion occurs when the particle is also positively charged and attraction occurs between opposite charges. For small ions, and in certain cases also proteins, this effect can be quantified by studying the ionic strength dependence of the reaction rate. The rate of association of oppositely charged ions will decrease with increasing ionic strength, the rate of association of the same charges will increase with increasing ionic strength, and when one of the particles is not charged there is no effect of the ionic strength.

[0055] The affinity of glucose isomerase for magnesium decreases with increasing ionic strength which is consistent with an overall negative charge of the glucose isomerase binding site. The binding of the cation may be altered by the introduction of a charged amino acids at the protein surface along the trajectory of the cation upon entrance of the active site. More specifically, this invention relates to the use of electrostatic forces to alter the association rate constant of the cation. Glucose isomerase may be engineered to increase the association rate for the cation by the addition of negative charge (or deletion of positive charge) near the active site channel, or to decrease the association rate for the cation by the addition of positive charge (or deletion of negative charge) near the active site channel. Since the off-rate is not expected to be affected substantially, an altered on-rate will result in an altered overall association constant of the cation. Since the loop regions situated at the C-termini of the  $\beta$ -barrel shape the active site entry, the possible mutation sites are searched in these regions. To avoid possible interference with barrel stability, substitutions in  $\beta$ -strands or  $\alpha$ -helices will not be considered. The following rational may be used:

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- Select all residues in the region between the C-terminal ends of the β-strands and the N-terminal ends of the αhelices.
- Reject from further consideration all residues where substitution leads to a decrease of the net positive charge in
  a sphere of 15 Angstroms radius around the metal ligands. Introduction of negative charges too close to the metal
  binding side will shift the pK<sub>a</sub> of the metal ligands to a higher pH, which will cancel out the effect of increased K<sub>ass</sub>
  at low pH.
- Compute for each of the remaining residues its accessible surface area in the context of the protein and using a
  probe of radius 1.4 Å. Reject residues that are buried in the sense that they have less than 10 Å<sup>2</sup> accessible surface
  area

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### STRUCTURAL INFORMATION

[0056] Information on the 3D structure of the enzyme (or enzyme:substrate or enzyme:inhibitor complex) is of great importance to be able to make predictions as to the mutations which can be introduced.

[0057] Structural data have been reported for glucose isomerase of <u>Streptomyces rubiginosus</u> (Carrell et al, J. Biol. Chem. 259 (1984) 3230-3236); Carrell et al. Proc. Natl. Acad. Sci USA 86, (1989) 440-4444) <u>Streptomyces olivochromogenus</u> (Farber et al, Protein Eng. 1, (1987) 467-475; Farber et al. Biochemistry <u>28</u> (1987) 7289-7297), <u>Arthrobacter</u> (Hendrick et al., J. Mol. Biol. 208 (1989) 127-157) and <u>Streptomyces albus</u> (Dauter et al FEBS Lett. 247, 1-8).

[0058] Although not all amino acid sequence data are available for these enzymes the 3D-structural homology with Actinoplanes missouriensis glucose isomerase is striking (see F. Rey et al., Proteins 4 (1988) 165-172). To show the general applicability of the method disclosed in this specification the genes for glucose isomerase originating from various species have been cloned and sequenced. The amino acid sequences of glucose isomerases as deduced from the genes of Streptomyces violaceoruber, Streptomyces murinus, Arthrobacter spec. and Streptomyces thermovulgaris are shown to be homologous. Published amino acid sequences for the glucose isomerases of Ampullariella sp. (Saari, ibid.) and Streptomyces violaceoniger (Nucl. Acids Res. 16 (1988) 9337), deduced from the nucleotide sequences of the respective genes, display a close homology to Actinoplanes missouriensis glucose isomerase. In addition, WO 89/01520 discloses that the amino acid sequence of Streptomyces rubiginosus glucose isomerase is homologous to Ampullariella sp. glucose isomerase.

[0059] Despite the absence of 3D structural data for most glucose isomerases, it can be concluded that all glucose isomerases from Actinomycetales have a similar tetrameric organisation.

[0060] In general, it can be assumed that where the overall homology is greater than 65%, preferably greater than 74% (minimal homology between <u>Actinoplanes missouriensis</u> and <u>Streptomyces</u> glucose isomerase, according to Amore and Hollenberg, Nucl. Acids Res. 17, 7515 (1989)), and more preferably greater than 85% and where the 3D structure is similar, amino acid replacements will lead to similar changes in pH optimum. Specifically one expects the glucose isomerases from species belonging to the order of the <u>Actinomycetales</u> to have such a high degree of similarity that the alteration of pH optimum due to amino acid replacements at the selected sites are similar. <u>Actinoplanes missouriensis</u> is the preferred source of glucose isomerase to mutate.

[0061] Figure 1 gives a schematic presentation of the active site of the glucose isomerase from Actinoplanes mis-

#### souriensis.

[0062] Figure 2 shows the aligned amino acid sequences of various glucose isomerases.

[0063] In the present specification both the three letter and the one letter code for amino acids is used (see e.g. Stryer, L. Biochemistry, p.13, 2nd ed, W.H. Freeman and Comp., NY, 1981).

#### **EXPERIMENTAL**

### Cloning and expression of the D-glucose isomerase gene

10 [0064] D-glucose isomerase (GI) is synonymously used for D-xylose isomerase ((D-xylose) ketol-isomerase, EC 5.3.1.5), an enzyme that converts D-xylose into D-xylose. The D-glucose isomerase from <u>Actinoplanes missouriensis</u> produced by engineered <u>E. coli</u> strains is designated as EcoAmi (DSM) GI. To distinguish the <u>Actinoplanes missouriensis</u> gene coding for GI from the analogous <u>E. coli xylA</u> gene, the former will be designated as <u>GI</u>.

[0065] Methods for manipulation of DNA molecules are described in Maniatis et al. (1982, Cold Sprong Harbor Laboratory) and Ausubel et al. (1987, Current Protocols in Molecular Biology, John Wiley & Sons Inc. New York). Cloning and DNA sequence determination of the glucose isomerase gene from <a href="Actinoplanes missouriensis">Actinoplanes missouriensis</a> DSM 43046 is described in EP-A-0351029.

The derived amino acid sequence of GI is numbered and compared with other glucose isomerases in Figure 2. In the following, the numbering of amino acids refers to Figure 2.

20 [0066] Wildtype and mutant Gi enzymes were produced in E. coli strain K514 grown as described in EP-A-0351029.

### Assay of the enzymatic activity of the expression product

[0067] The enzymatic activity of glucose isomerase was assayed as described below (1 unit of enzymatic activity produces 1.0 micromole of product -D-xylulose or D-fructose-per minute; therefore, specific activity -spa- is expressed as units per mg of GI enzymes).

[0068] GI activity can be assayed directly by measuring the increase in absorbance at 278 nm of xylulose produced at 35°C by isomerisation of xylose by glucose isomerases. This assay was performed in 50 mM triethanolamine buffer, pH 7.5, containing 10mM MgSO<sub>4</sub>, in the presence of 0.1 M xylose. Glucose isomerase final concentration in the assay was  $\pm$  0.01 mg/ml, and precisely determined, prior to dilution in the enzymatic assay mixture, by absorption spectroscopy using an extinction coefficient of 1.08 at 278 nm for a solution of enzyme of 1.0 mg/ml.

[0069] The specific activity was determined in the D-Sorbitol Dehydrogenase Coupled Assay, enzymatic determination of D-xylulose was performed at 35°C as previously described (Kersters-Hilderson et al., Enzyme Microb. Technol. 9 (1987) 145) in 50mM triethanolamine, pH 7.5, 10mM MgSO<sub>4</sub>, and 0.1 M xylose, in the presence of  $\pm$  2 x 10-8 M D-sorbitol dehydrogenase (L-iditol: NAD oxidoreductase, EC 1.1.14), and 0.15 nM NADH, except that the incubation buffer also included 1mM ethylenebis(oxyethylenenitrilo)tetraacetic acid (EGTA). Glucose isomerase final concentration in this assay was  $\pm$  2.5 x 10<sup>3</sup> mg/ml, and precisely determined as described above.

[0070] With glucose as a substrate GI activity can be assayed by the measurement of D-fructose produced during the isomerization reaction using the cysteine-carbazole method (CCM) which is based on the reaction of ketosugars with carbazole in acids to yield a purple product (Dische and Borenfreund, J. Biol. Chem. 192 (1951) 583).

### **EXAMPLE 1**

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### The pH dependence of glucose isomerase activity

[0071] In order to determine the pH-activity profile of wild-type and mutant glucose isomerase, the activity was measured as a function of pH (5.2-8.0) in the presence of 10 mM MgSO<sub>4</sub> and 200 mM xylose (using the direct assay method). For mutants with very low activity the coupled sorbitol dehydrogenase assay system was used between pH 5.8 and 8.4. Care was taken that the sorbitol dehydrogenase reaction did not become rate limiting at extreme pH values.

[0072] The pH-activity profile of glucose isomerase (i.e. the recombinant wildtype enzyme from <u>Actinoplanus missouriensis</u>) in the presence of 200 mM xylose and different activating cations is shown in Figure 3. It reveals a decrease in activity at both acidic pH below 7.0 and at alkaline pH beyond 8.0.

[0073] The appropriate steady-state kinetic mechanism for glucose isomerase involves the rapid formation of an enzyme-metal-sugar complex which is converted to the product in a rate limiting step (so called rapid equilibrium, random ordered mechanism - see Figure 4). Equilibrium and transient kinetic fluorescence measurements (stopped flow) indicate the presence of two metal ion binding sites. In the stopped flow experiment the metal ions bind consecutively. The high affinity metal plays a role in maintaining an active conformation and is therefore called the 'conformational' site. The second metal binding site accommodates the activating cation, therefore this site is usually indicated

as the 'catalytic' site. The reaction scheme which is shown in Figure 4 appears to be adequate to analyze and compare steady-state and stopped flow experiments. In principle steady-state results do not distinguish between the two metal binding sites, but it is assumed that the main effect comes from the catalytic metal binding.

[0074] Analysis of the initial rate (v) of xylose conversion as a function of the xylose and magnesium concentrations allows to determine four parameters: the maximal velocity, the equilibrium constants for magnesium binding to the free enzyme ( $K_1$ ), the enzyme-sugar complex ( $K_4$ ), and the equilibrium constants of xylose binding to the free enzyme ( $K_2$ ) and the enzyme-magnesium complex ( $K_3$ ).

$$\frac{V_{max}}{v} = 1 + \frac{1}{K_3^*[S]} + \frac{1}{K_4^*[M]} + \frac{1}{K_3^*K_1^*[S]^*[M]}$$

where [M] and [S] represent the concentration of the metal ion and xylose respectively.

[0075] By systematic variation of the magnesium ion and the xylose concentrations it was possible to obtain values for the maximal velocity and for all four equilibrium constants. Figure 5 shows the pH dependance of these parameters. [0076] Comparison of the results of Figures 3 and 5, shows that the acidic side of the pH profile is completely determined by the metal ion binding.

The data of Figure 5 are not sufficiently accurate to calculate the number of ionizations involved. The slope of the plots of  $logK_{1,4}$  vs pH (slope > 1) indicates however, that more than one ionization may be involved. The similarity in pH dependence of  $K_1$  and  $K_4$  indicates that the same ionizing groups are important for both these processes (involving the same site).

#### **EXAMPLE 2**

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Selection of amino acid residues in glucose isomerase of which substitution will alter the pK<sub>a</sub> values of the metal binding, ionizable amino acids.

[0077] In the case of glucose isomerase, the criteria for the selection of possible positions for substitution, as outlined in the detailed description of this invention, were applied using the aligned sequences from different sources (Figure 2) and the highly refined structure of <a href="Actinoplanes missouriensis">Actinoplanes missouriensis</a> glucose isomerase in complex with the inhibitor xylitol (see Figure 1 and "Structural Information" in the "Detailed description of the invention"). The highly refined structure with a resolution of 2.2 Angstroms reveals the position of the inhibitor and two metal binding sites. A schematic representation of the active site of glucose isomerase of Actinoplanes missouriensis is given in Figure 1.

[0078] From the glucose isomerase structure complexed with cobalt and xylitol those residues were selected where substitution into a more positively charged amino acid residue leads to a net increase in positive charge within a 15 Angstrom radius around the target ionizable groups. In the case of glucose isomerase the targets are the ionizable goups that are involved in the coordination of the metal ions required for activity. These target ionizable groups imply the carboxylgroups of Glu181, Glu217, Asp245, Asp292 and the NE of His220.

[0079] After application of criterion I, 80 possible mutation sites were retained. These sites are summarized below:

Ala5, Phe11, Leu15, Trp20, Gln21, Ala25, Phe26, Asp28, Ala29, Gly47, Tyr49, Thr52, Phe53, His54, Asp56, Asp57, Phe61, Ile85, Met88, Phe94, Thr95, Phe104, Gln122, Thr133, Leu134, Val135, Ala143, Tyr145, Tyr158, Asn163, Ser169, Glu181, Asn185, Glu186, Gly189, Ile191, Pro194, His198, Gln204, Leu211, Phe212, Asn215, Glu217, Thr218, His220, Glu221, Gln222, Ser224, Asn225, Leu226, Phe228, Thr229, Gly231, Leu236, His238, His243, Asp245, Asn247, His250, Phe254, Asp255, Gln256, Asp257, Leu258, Val259, Phe260, His262, Leu271, Tyr285, Asp286, His290, Asp292, Tyr293, Thr298, Glu299, Trp305, Ala310, Met314, Val380, Asn383.

[0080] After discarding the catalytic residues and the strictly conserved residues (criterion II) the following 62 residues are left:

Ala5, Leu15, Gin21, Ala25, Phe26, Asp28, Ala29, Gly47, Tyr49, Thr52, Asp56, Phe61, Ile85, Met88, Thr95, Phe104, Gln122, Thr133, Leu134, Ala143, Tyr145, Tyr158, Asn163, Ser169, Asn185, Gly189, Ile191, Pro194, His198, Gln204, Leu211, Phe212, Thr218, Gln222, Ser224, Asn225, Leu226, Phe228, Thr229, Gly231, Leu236, His238, His243, His250, Phe254, Gln256, Asp257, Leu258, Val259, His262, Leu271, Tyr285, Asp286, His290, Tyr293, Thr298, Glu299, Trp305, Ala310, Met314, Val380, Asn383.

[0081] Subsequently a priority was attributed to each of these 62 possible mutation sites according to criterion III. The following 24 sites were attributed as having high priority for mutagenesis: Ala25, Gly47, Tyr49, Thr52, Phe61,

Ile85, Thr95, Gln122, Thr133, Tyr145, Tyr158, Gly189, Ile191, Gln204, Thr218, Gln222, Leu226, Thr229, Gly231, Leu236, Gln256, Leu258, Tyr293 and Val380.

[0082] Mutation of either one or several of the 80 selected amino acid residues into positively charged residues will result in a decrease of the  $pK_a$  of metal binding of glucose isomerase. This may result in a corresponding shift of the pH-activity profile towards lower pH.

[0083] Correspondingly, mutation of either one or several of the 80 selected amino acid residues into negatively charged residues will result in an increase of the pK<sub>a</sub> of metal binding of glucose isomerase. This may result in a corresponding shift of the pH-activity profile towards higher pH.

#### 10 REFERENCE EXAMPLE A

The effect of mutating Lys 294 on the pH-activity profile

[0084] At position 294 in glucose isomerase a positive charge is located about 8 Angstroms from the highest occupied metal binding site in the glucose isomerase-xylitol complex (395 in Figure 1). A mutation at this site was made, replacing the lysine for an arginine. This mutation conserves the positive charge at position 294. Consistent with this conservation of the positive charge is the observation that the pH-activity profile of this mutant is similar to that of the wildtype enzyme.

[0085] However, when at position 294 the positive charge is removed by replacing lysine 294 by a glutamine we observed a shift of the pH-activity profile towards the alkaline site by approximately 0.5 pH units. The pH-activity profiles for K294R and K294Q are shown in Figure 6.

[0086] This example illustrates that it is possible to manipulate the  $pK_a$  of one or more functional groups, in the active site of glucose isomerase, by changing the net charge of the protein around the active site.

#### REFERENCE EXAMPLE B

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The effect of mutating Glu186 and replacing magnesium ions by manganese ions on the pH-activity profile

[0087] In the mutant E186Q a negative charge is replaced by a neutral one which gives rise to a net increase in positive charge within a 15 Angstroms radius around the metal ligands. The pH-activity profile of the E186Q mutant in the presence of magnesium is shown in Figure 7. The alkaline flank of the pH profile is shifted significantly toward lower pH. In the presence of magnese instead of magnesium the pH-activity profile of E186Q is shifted to a lower pH and at its optimum pH its activity is higher than for the wildtype (Figure 8).

[0088] For applications where metal ions other than magnesium can be used the combination E186Q with manganese at low pH is an interesting option.

[0089] We also performed the mutation E186D which is conservative with respect to charge. The pH-activity profile of this mutant is shown in Figures 7 and 8. The pH dependence of the activity for the E186D mutant is not significantly different from that of the wildtype enzyme. Removal of the negative charge at position 186 did shift the pH activity profile to a more acidic pH. This example emphasizes that the rationale of the mutation E186Q holds.

## 40 REFERENCE EXAMPLE C

The effect of replacing Asp255 on the pH-activity profile

[0090] Substitution of the negatively charged aspartic acid at position 255, which is in fact a metal binding ligand in glucose isomerase, by a neutral asparagine, gives rise to a shift of the pH-activity profile towards lower pH in the presence of manganese. The pH optimum shifts about 2 pH units towards more acidic pH.

[0091] The pH-activity profile is given in Figure 9.

### **EXAMPLE 3**

Glucose isomerases with an altered pH-activity profile

[0092] Mutants of glucose isomerase which were created according to the methods as outlined in the detailed description of the invention, were tested for their pH-activity relation under conditions which are indicated in the Figures (10-20). The pH-activity profile of a mutant is the result of the effect of the mutation on the  $pK_a$  on the one hand, and the effect on the  $K_{ass}$  on the other hand.

[0093] The results for the following mutants are given in the Figures:

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F254K (Fig.10), F94R (Fig.11), F61K (Fig.12), A25K (Fig.13), D57N (Fig.14), L258K (Fig.15), Q204K (Fig.16), R23Q (Fig.17), H54N (Fig.18), H290N (Fig.19), T95D (Fig.20).

[0094] For the mutants in which a positive charge (F254K, F94R, F61K, A25K, L258K, Q204K) was introduced or a negative charge neutralized (D57N), it can be seen that the acidic-side of the pH-activity profile is shifted towards lower pH.

[0095] For mutants in which a negative charge was introduced (T95D) or a positive charge was neutralized (R23Q, H54N, H290N, it can be seen that the pH-activity profile is shifted towards higher pH.

[0096] Both of these observations are in agreement with the model as presented in the detailed description of the invention.

[0097] However, it should be noted that mutants in which a conserved amino acid has been replaced (F94R, D57N, H54N) give a drastic decrease in specific activity on xylose. At position 254 in the sequence alignment (Fig. 2) only hydrophobic amino acids are found. Introduction of a charged amino acid (F254K) at this position also leads to a drastic decrease in specific activity. Thus, it can be concluded that although (semi)conserved amino acid positions can be used to alter charges in order to modify the pH-activity profile they are not preferred sites.

### **EXAMPLE 4**

Stabilization of mutants with an altered pH optimum to obtain better performance under application conditions

[0098] The mutants H290N and F61K give the expected shift in the pH-activity profile as described in Example 6. Of these mutants H290N was immobilized as described in EP-A-351029 (Example 7). Application testing of the wildtype and this mutant glucose isomerase was performed as described in the same application (Example 8). The stability is indicated by the first order decay constant ( $K_d$ , the lower the decay constant the more stable the enzyme). Table 2 gives the  $K_d$  values for the wild-type and mutant glucose isomerases.

Table 2

| Decay constants for wildtype and mutant glucose isomerase, immobilized on Lewatit |   |  |  |  |
|---|---|--|--|--|
|   | K <sub>d</sub> (x 10 <sup>6</sup> sec <sup>-1</sup> ) |  |  |  |
| Wildtype  | 2.5   |  |  |  |
| H290N   | 3.1   |  |  |  |
| K253R   | 0.7   |  |  |  |
| R290NK253R  | 1.6   |  |  |  |
| F61KK253R   | 1.4   |  |  |  |

[0099] As can be seen in Table 2, H290N is destabilized as compared with the wildtype glucose isomerase. K253R was found to stabilize the wildtype glucose isomerase by a factor larger than three. The pH-activity profile of the K253R mutant is identical with that of the wildtype enzyme.

The combination of the pH mutant H290N with the stability mutant K253R shows that pH mutants can be stabilized by introducing mutations that have been shown to stabilize the wildtype enzyme.

[0100] In addition Table 2 shows that pH mutant F61K is stabilized with respect to the wiltype enzyme after introduction of K253R.

[0101] The acidic shift of the pH-activity profile of F61K is maintained in the double mutant (Fig. 21). This shows that mutations which improve different properties of an enzyme can be combined in a new mutant which harbours the improved properties of the individual mutations (an improved pH optimum and an improved stability in this case).

[0102] It is to be understood that the above mentioned examples are meant to demonstrate the concept of the invention which is defined in the claims. It should be clear that combinations of the above mentioned mutations with other mutations leading to altered characteristics, e.g. thermostability, metal binding or substrate specificity, are also encompassed by the subject invention.

#### Claims

A method for altering the pH-activity profile of a wild-type glucose isomerase enzyme such that said profile, or the
acidic part thereof, is shifted towards a lower pH, comprising the substitution of a negatively charged amino acid
residue (Asp or Glu) by a neutral one or by a positively charged one, or the substitution of a neutral residue by a

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positively charged one (Arg or Lys), wherein said substitution is at a position selected from the group consisting of:

Ala25, Phe61, Gln204, and Leu258;

- wherein said residues of the glucose isomerase are from <u>Actinoplanes missouriensis</u> whose sequence is shown in Figure 2, or the corresponding position in a homologous glucose isomerase.
  - A method according to claim 1 wherein the wild-type glucose isomerase is obtained from a microorganism of the order <u>Actinomycetales</u>.
  - 3. A method according to claim 2, wherein the wild-type glucose isomerase is obtained from <u>Actinoplanes missouriensis</u>.
- A method according to any one of the preceding claims wherein the glucose isomerase contains at least one of the following amino acid replacements: A25K, F61K, Q204K, and L258K.
  - A method according to claim 1 wherein the glucose isomerase is the <u>Actinoplanes missouriensis</u> isomerase of Figure 2 and contains the replacement F61K+K253R.
- 20 6. A process for obtaining a mutant glucose isomerase molecule which comprises:
  - (a) obtaining a DNA sequence encoding a glucose isomerase;
  - (b) mutating this sequence at selected nucleotide positions to provide an enzyme altered at any one of positions Ala25, Phe61, Gln204, and Leu258 by a substitution of a negatively charged amino acid residue (Asp of Glu) by a neutral one or by a positively charged one, or the substitution of a neutral residue by a positively charged one (Arg or Lys),

wherein said residues of the glucose isomerase are from <u>Actinoplanes missouriensis</u> whose sequence is shown in Figure 2, or the corresponding position in a homologous glucose isomerase;

- (c) cloning the altered sequence into an expression vector in such a way that the DNA sequence can be expressed;
- (d) transforming a host organism or cell with said vector;
- (e) culturing said host organism; and,
- (f) isolating the altered glucose isomerase.
- 7. The process of claim 6 wherein the wild-type glucose isomerase is obtained from a microorganism of the order Actinomycetales.
  - 8. The process of claim 7, wherein the wild-type glucose isomerase is obtained from Actinoplanes missouriensis.
- 9. The process of any one of claims 6 to 8 wherein the glucose isomerase contains at least one of the following amino acid replacements: A25K, F61K, Q204K, and L258K.
  - 10. The process of claim 6 wherein the glucose isomerase is the <u>Actinoplanes missouriensis</u> isomerase of Figure 2 and contains the replacement F61K+K253R.
  - 11. The process of any one of claims 6 to 10 which further comprises the use of the mutant glucose isomerase in the production of a fructose syrup.

#### 50 Patentansprüche

1. Verfahren zur Änderung des pH-Aktivitätsprofils eines Wildtyp-Glucoseisomerase-Enzyms derart, dass das Profil oder der saure Teil desselben in Richtung auf einen niedrigeren pH verschoben wird, umfassend die Substitution eines negativ geladenen Aminosäure-Restes (Asp oder Glu) durch einen neutralen oder durch einen positiv geladenen oder die Substitution eines neutralen Restes durch einen positiv geladenen (Arg oder Lys), wobei die Substitution an einer Position stattfindet, die ausgewählt ist aus der Gruppe bestehend aus:

Ala25, Phe61, Gln204 und Leu258;

wobei die Reste der Glucoseisomerase aus Actinoplanes missouriensis, deren Sequenz in Figur 2 gezeigt ist, oder der entsprechenden Position In einer homologen Glucoseisomerase sind.

- Verfahren nach Anspruch 1, in dem die Wildtyp-Glucoseisomerase aus einem Mikroorganismus der Ordnung Actinomycetales erhalten wird.
  - 3. Verfahren nach Anspruch 2, in dem die Wildtyp-Glucoseisomerase aus Actinoplanes missouriensis erhalten wird.
- Verfahren nach irgendeinem der vorangehenden Ansprüche, in dem die Glucoseisomerase mindestens eine der folgenden Aminosäure-Ersetzungen enthält: A25K, F61K, Q204K und L258K.
  - 5. Verfahren nach Anspruch 1, in dem die Glucoseisomerase die Actinoplanes missouriensis-Isomerase von Figur 2 ist und den Ersatz F61K+K253R enthält.
- 15 6. Verfahren zum Erhalten eines mutierten Glucoseisomerase-Moleküls, welches umfasst:
  - (a) Erhalten einer DNA-Sequenz, die eine Glucoseisomerase codiert;
  - (b) Mutieren dieser Sequenz an ausgewählten Nukleotid-Positionen, um ein Enzym bereitzustellen, das an irgendeiner der Positionen Ala25, Phe61, Gln204 und Leu258 durch eine Substitution eines negativ geladenen Aminosäurerestes (Asp oder Glu) durch einen neutralen oder einen positiv geladenen oder die Substitution eines neutralen Restes durch einen positiv geladenen (Arg oder Lys) geändert ist, wobei die Reste der Glucoseisomerase aus Actinoplanes missouriensis, deren Sequenz in Figur 2 gezeigt ist, oder die entsprechende Position in einer homologen Glucoseisomerase sind;
  - (c) Klonieren der geänderten Sequenz in einen Expressionsvektor auf solche Weise, dass die DNA-Sequenz exprimiert werden kann;
  - (d) Transformieren eines Wirtsorganismus oder einer Wirtszelle mit dem Vektor;
  - (e) Kultivieren des Wirtsorganismus; und
  - (f) Isolieren der geänderten Glucoseisomerase.
- Verfahren nach Anspruch 6, in dem die Wildtyp-Glucoseisomerase aus einem Mikroorganismus der Ordnung Actinomycetales erhalten wird.
  - 8. Verfahren nach Anspruch 7, in dem die Wildtyp-Glucoseisomerase aus Actinoplanes missouriensis erhalten wird.
- Verfahren nach irgendeinem der Ansprüche 6 bis 8, in dem die Glucoselsomerase mindestens eine der folgenden Aminosäure-Ersetzungen enthält: A25K, F61K, Q204K und L258 K.
  - Verfahren nach Anspruch 6, in dem die Glucoseisomerase die Actinoplanes missouriensis-Isomerase von Figur 2 ist und den Ersatz F61K+K253R enthält.
  - Verfahren nach Irgendeinem der Ansprüche 6 bis 10, das weiter die Verwendung der mutierten Glucoseisomerase bei der Herstellung eines Fructosesirups umfasst.

## 45 Revendications

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1. Procédé de modification du profil d'activité pH d'une enzyme isomérase de glucose de type sauvage de telle sorte que ledit profil, ou la partie acide de celui-ci, est décalé vers un pH inférieur, ledite procédé comprenant la substitution d'un résidu aminoacide chargé négativement (Asp ou Glu) par un résidu neutre ou par un résidu chargé positivement, ou la substitution d'un résidu neutre ou par un résidu chargé positivement (Arg ou Lys), ladite substitution se faisant à une position choisie dans le groupe constitué par :

Ala25, Phe61, Gln204, et Leu258;

- lesdits résidus de l'isomérase de glucose provenant de <u>Actinoplanes missouriensis</u> dont la séquence est représentée dans la figure 2, ou la position correspondante dans une isomérase de glucose homologue.
- 2. Procédé selon la revendication 1, dans lequel l'isomérase de glucose de type sauvage est obtenue à partir d'un

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#### EP 0 436 502 B1

micro-organisme de l'ordre Actinomycetales

- Procédé selon la revendication 2, dans lequel l'isomérase de glucose de type sauvage est obtenue à partir de <u>Actinoplanes missouriensis.</u>
- Procédé selon l'une quelconque des revendications précédentes, dans lequel l'isomérase de glucose contient au moins un des remplacements aminoacides suivants : A25K, F61K, Q204K, et L258K.
- Procédé selon la revendication 1, dans lequel l'isomérase de glucose est l'isomérase de <u>Actinoplanes missourien-sis</u> de la figure 2 et contient le remplacement F61K+K253R.
  - .6. Processus d'obtention d'une molécule d'isomérase de glucose mutante, lequel processus comprend :
    - (a) l'obtention d'une séquence d'ADN codant une isomérase de glucose ;
    - (b) la mutation de cette séquence à des positions de nucléotides sélectionnées pour fournir une enzyme modifiée en une position quelconque des positions Ala25, Phe61, Gln204, et Leu258 par une substitut ion d'un résidu aminoacide chargé négativement (Asp ou Glu) par un résidu neutre ou par un résidu chargé positivement, ou la substitution d'un résidu neutre par un résidu chargé positivement (Arg ou Lys),

lesdits résidus de l'isomérase de glucose provenant de <u>Actinoplanes missouriensis</u> dont la séquence est représentée dans la figure 2, ou la position correspondante dans une isomérase de glucose homologue ; (c) le clonage de la séquence modifiée en un vecteur d'expression de telle manière que la séquence d'ADN peut être exprimée ;

- (d) la transformation d'un organisme hôte ou d'une cellule hôte avec ledit vecteur ;
- (e) la culture dudit organisme hôte; et,
- (f) l'isolation de l'isomérase de glucose modifiée.
- Processus selon la revendication 6, dans lequel l'isomérase de glucose de type sauvage est obtenue à partir d' un micro-organisme de l'ordre Actinomycetales.
- 8. Processus selon la revendication 7, dans lequel l'isomérase de glucose de type sauvage est obtenue à partir de Actinoplanes missouriensis.
  - 9. Processus selon l'une quelconque des revendications 6 à 8, dans lequel l'isomérase de glucose contient au moins un remplacement parmi les remplacements aminoacide suivants : A25K, F61K, Q204K, et L258K.
  - Processus selon la revendication 6, dans lequel l'isomérase de glucose est l'isomérase de <u>Actinoplanes</u> <u>missouriensis</u> de la figure 2 et contient le remplacement F61K+K253R.
- 11. Processus selon l'une quelconque des revendications 6 à 10 qui comprend en outre l'utilisation de l'isomérase de glucose mutante dans la production d'un sirop contenant du fructose.

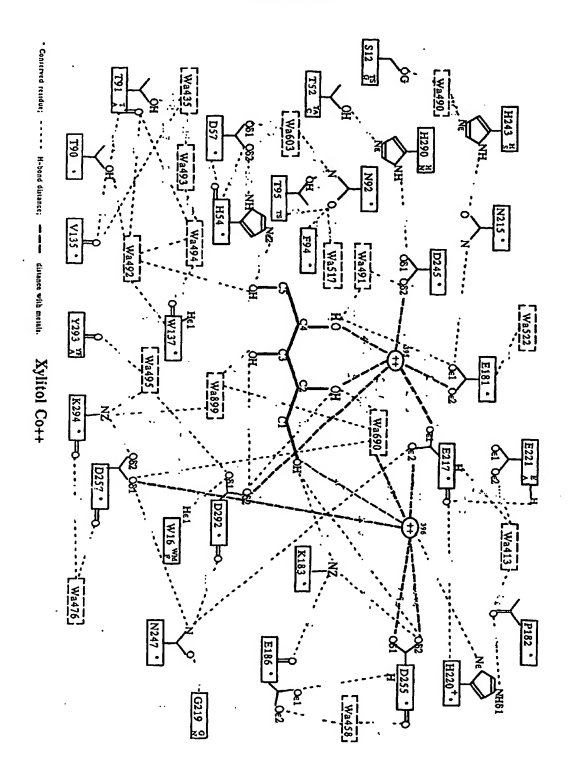


FIGURE 1

| M SVQAIRED AE SECLUTVOWQ AR. DAFCDAIR H SLOATPDD AT SECLUTVOWQ AR. DAFCDAIR H NYQPIPED AT THE LUTVOWQ CR. DFFCDAIR M SYQPIPED AT THE LUTVOWQ CR. DFFCDAIR M SYQPIPED RE SECLUTVOWQ CR. DFFCDAIR M SYQPIPED RE SECLUTVOWQ CR. DFFCDAIR M SYQPIPADHE TWO LATVOWY CR. DFFCDAIR MANYPOQLDRVRYEGSKSSNPLAFRHYNPDELVLOKRUZEHLRFSIAYWH IFTADGID WF DR. TUQRRFWD MY.KGM. DLAR MANYPUDIAPIKYEGILIKNNFAFRHYNPDELVLOKRUZEHLRFALYMH IFTWDGAD WF GYG AFNRPWQ QPGEAL. ALAK | TAL DPV<br>PVL DPI<br>QAL DPA<br>PAL DPV<br>RPL DPV<br>KNL DPV<br>ARV EAA<br>RKA DVA |  |
|---|--|--|
| <b>пилипили</b>   | 11 11 11 11 11 11 11 11 11 11 11 11 11   | 1111 1111 1111 1111 1111 1111 1111 1111 1111         |
| ABS<br>SAS<br>SBU<br>SBU<br>Art<br>Art<br>Art<br>BSU  | Amb<br>Amp<br>Svi<br>Smu<br>Sth<br>Art<br>Art<br>Eco                                 | Ami<br>Ami<br>Swi<br>Smu<br>Sth<br>Art<br>Art<br>Eco |

FIGURE 2A

| IG DLLNAFEL IG DLLNAFEL AG DLRAAFWL AG DLRAAFWL AG DLRAAFWL AG DLRAAFWL AG DLRAAFWL AG VEENALWA NS VEENALWA TL VIDITLAM  | ELKTPTLNPG ELRTPTLNPG ELRAPTAA. QLAQPTAA. QLAQPTAA. ELGETTLNAG F.TEGIGLEI W.NSELGQQI   |   |   |
|--|--|---|---|
| PKFDQDLVFG PKFDQDLVFG IKYDQDLNFG IKYDQDLNFG IKYDQDLNFG IKYDQDLNFG IKYDQDLNFG ICWDTDE.FP LCWDTDE.FP LCWDTDE.FP  | QEALAASKVA<br>QAALAESKVD<br>QEALAAARLD<br>QEALRAARLD<br>QEALRASRLD<br>QEALKTSCVF<br>EDVIQHRYRS<br>DKRIAQRYSG<br>SDIVDERYSS   |   |   |
| HIDE NGONG . HIDE NGONG . HIDE NGOSG . HIDE NGOSG . HIDE NGOSG . HIDE NGONG . SYDA NGONG L   | AKAFRA DPEV<br>AKAFRA DPEV<br>AAAFRA DPEV<br>SAAFRA DPEV<br>ALAFRA DPEV<br>HKLI.E DRVF<br>ARMI.E DGEL  |   |   |
| DALWHK K. F. P.  | IRMYLLLKER IRMYLLLKER MRNYLILKDR MRNYLILKDR MSKYLLLKER DAFARGLKVA DTWALALKIA   | CAR<br>CAR<br>CAR<br>CARG   | GSR   |
| INL NFTGG IANGEL | DGVWESAKAN<br>DGVWASAKCO<br>DGVWASAAGC<br>DGVWASAAGC<br>DGVWASAAGC<br>DLVYAHIAGM<br>DLFYGHIGAM   | . Lnqlaiehll<br>. Lnqlaidhll<br>. Ldqlaidhll<br>. Ldqlaidhll  | .Lnqlaiehll<br>ilnq<br>Lvnhylfdk<br>Vlndylv   |
| LIP TAGHAIA FVQ ELERPE D'ECENFET CH. CASNL NFTGG IAQALWHKK METHIDE NCCHG PREDGDLYFG HG DLLNAFSL LLP TAGHAIA FVQ ELERPE PRIMERICAMSNL NFTGG IAQALWHKK METHIDE NCGSG IKYDQDLYFG HG DLLNAFSL LLP TVGHALA FIE RLERPE WONNEEV CHE QMACL NFPHG IAQALWACK LEHIDE NCGSG IKYDQDLRFG AG DLRAAFWL LLP TVGHALA FIE RLERPE WONNEEV CHE QMACL NFPHG IAQALWACK LEHIDE NCGSG IKYDQDLRFG AG DLRAAFWL LLP TVGHALA FIE RLERPE WONNEEV CHE QMACL NFPHG IAQALWACK EHIDE NCGSG IKYDQDLRFG AG DLRAAFWL LLP TVGHCLA FIE QLEHGDE VCENWEEV CHE QMACL NFPHG IAQALWACK EHIDE NCGSG IKYDQDLYFG HG DLISAFFT YDT DAATTIA FIK QYGLDN FRINKEEANHA ILAGH FFEHE LRMARVHG LE GSYDA NCGHF LLGWDTDE.FP TD LYSTILAM YDYD DAATVYG FIK GFGLEK FIKENTEANHA ILAGH FFEHE IATAIALG ESYDA NGCHF LLGWDTDE.FP TD LYSTILAM YDD DSATALA FLQKYDLDK ENNAGER NHA YLAGH FFEHE IATAIALG ESYDA NGCHF LLGWDTDE.FP TL VIDITIAM   | VDLLE NG. PDG APAXDGP RHF D YKPSRT. E DY DGVWESAKAN IRMYLLLKER AKAFRA DPEV GALLASKVA ELKIPILNPG VDLLE NG. PDG GPAYDGP RHF D YKPSRT. E DF DGVWESAKDN IRMYLLLKER AKAFRA DPEV GALLASKUD ELRIFILNPG VDLLE RAGYAGP RHF D FKPPRT. E DF DGVWASAAGG WRNYLLIKDR AAFRA DPEV GEALRARLD GLAGFTAA VDLLE TAGYDGP RHF D FKPPRT. E DF DGVWASAAGG WRNYLLIKDR AAFRA DPEV GEALRARLD GLAGFTAA VDLLE SSGYDGP RHF D FKPPRT. E DL DGVWASAAGG WRNYLLIKER SAAFRA DPEV GEALRASRLD GLAGFTAA VDLLE NGFPNG GPKYTGP RHF D FKPPRTE DL DGVWASAAGG WRNYLLIKER ALAFRA DPEV GEALRASRLD GLAGFTAA VDLLE NGFPNG GPKYTGP RHF D YKPSRTD GY DGVWDSAKAN WSWYLLIKER ALAFRA DPEV GEAWKTSGYF ELGETTLNAG VDLLE NGFPNG GPKYTGP RHF D YKPSRTD GY DLYVAHTAGW DAFARCLKVA HKLI.E DRVF EDVIGHRYRS F.TEGIGLEI YEILK AGGF.TTGGINF D AKVRRSSFE PD DLYVAHTAGW DTWARALKTA ARWI.E DOEL DKRIAGRYSG W.NSELGQOI YEILK AGGF.TTGGINF D AKVRRGSTD KY DLFYGHIGAW DTWARALKGA AAII.E DKFL SDIVDERYSS YRNTEVGGSI | ECYAELLADR SAFED.YDAD AVGAKGFGFVK .LNGLAIEHLL GAR ETYADLLADR SAFED.YDAD AVGAKGYGFVK .LNGLAIDHLL GAR DGLAALLADR SAYDT.FDVD AAAARGWAFEH .LDGLAWDHLL GAR DGLDALLADR AAFED.FDVD AAAARGWAFEH .LDGLAWDHLL GAR | AAAERNFAFIR .LNQLAIEHLL GSR<br>NESGRGERLKP ILNQ<br>HQSGRGEQLEN LVNHYLFDK<br>LDSNHLEYIKS VLNDYLV     |
| VQ ELERPE L F<br>IE RLERPE L Y<br>IE RLERPE L Y<br>IE RLERPE L Y<br>IE QLEHGD I W<br>LK QYGLDN H F<br>LK QFGLEK F T  | APAYDGP RUF GPAYDGP RUF CONTROL Y AGG RUF  | AFED.Y. DAD<br>AYDI.F. DVD<br>AYED.F. DVD   | SFAG.FDAE<br>QYALNNK.TIK<br>KYAQEHHLSPV<br>AFALEYGDDIE  |
| LLP TAGHAIA F LLP TAGHAIA F LLP TVGHALA F LLP TVGHALA F FLP TVGHALA F  | VDLLE NG. PDG<br>VDLLE RAG<br>VDLLE RAG<br>VDLLE SSG<br>VDLLE SSG<br>VDLLE NGFPNG<br>VDLLE NGFPNG<br>VDLLE NGFPNG<br>VDLLE NGFPNG<br>VDLLE NGFPNG<br>VDLLE NGFPNG  | ECYAELLADR SAFED.Y. DAD ETYADLLADR SAFED.Y. DAD DGLAALLADR SAYDT.F. DVD DGLDALLADR AAFED.F. DVD   | DGLQ ESAADLMNDS ASFAG.FDAE TEGRANFHTL EQYALNNK.TIK LKGQMSLADL AXYAQEHHLSPV FNGTATFFSI. AAFALEYGDDLE |
| 191<br>191<br>191<br>191<br>191<br>248<br>242  | 271<br>271<br>271<br>271<br>271<br>271<br>271<br>3228<br>322   | Ani 351<br>Anp 351<br>Svi 344 '<br>Sau 344 '  | Sth 348 Art 352 Bsu 404 Eco 380   |
| ABB<br>Soft<br>Soft<br>Soft<br>Soft<br>Soft<br>Soft<br>Soft<br>Soft  | LX St  | SAS   | N K K K   |

PIGURE 2

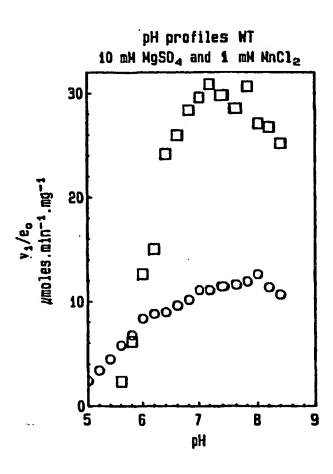
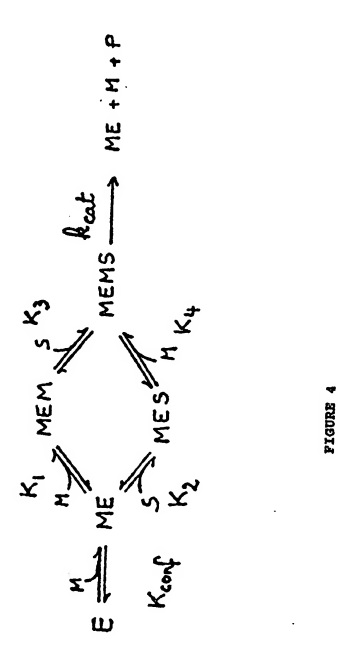
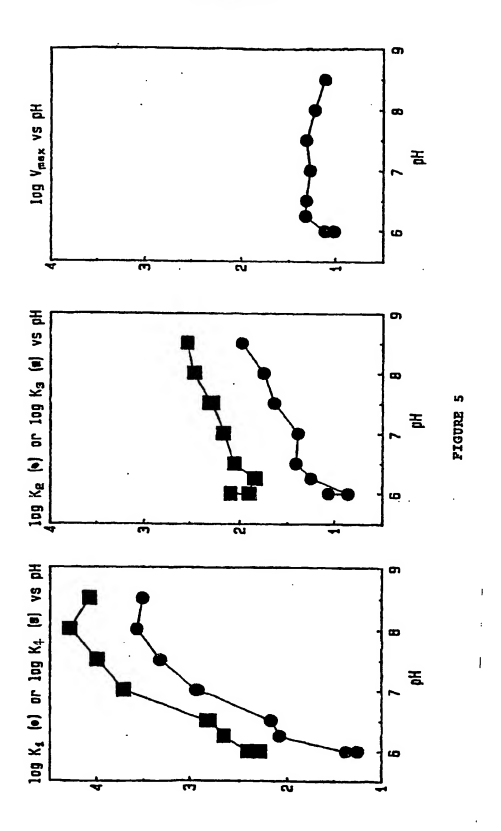


FIGURE 3





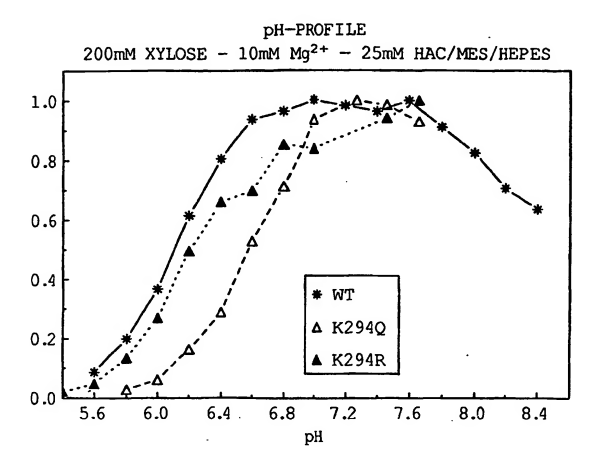


FIGURE 6

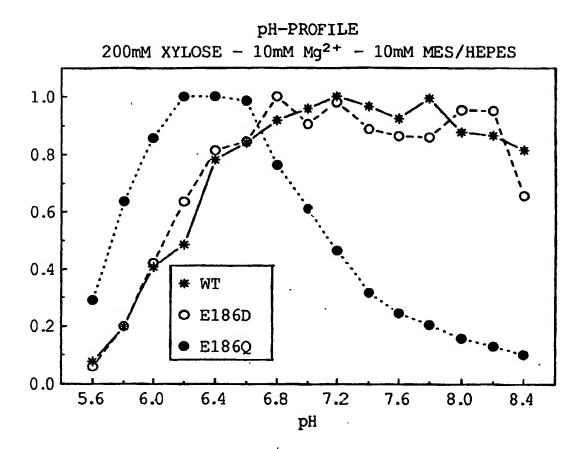
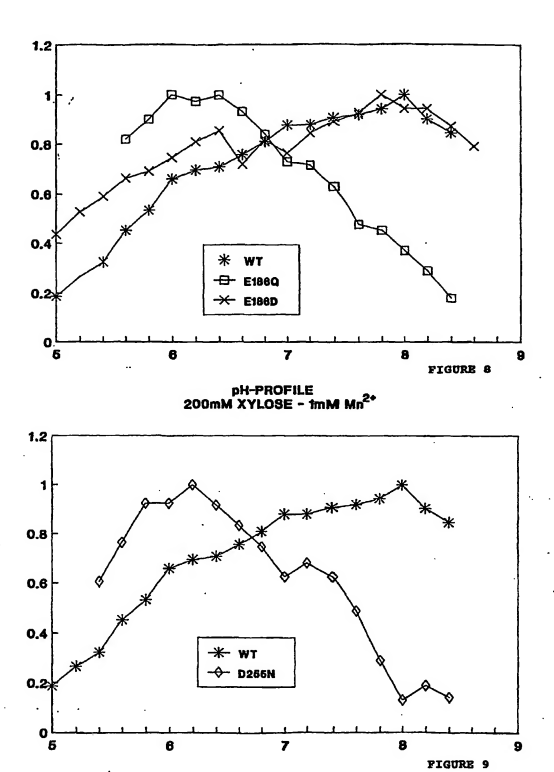


FIGURE 7



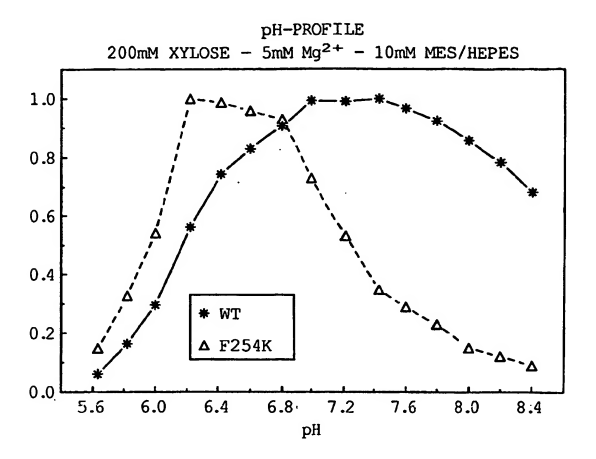


FIGURE 10

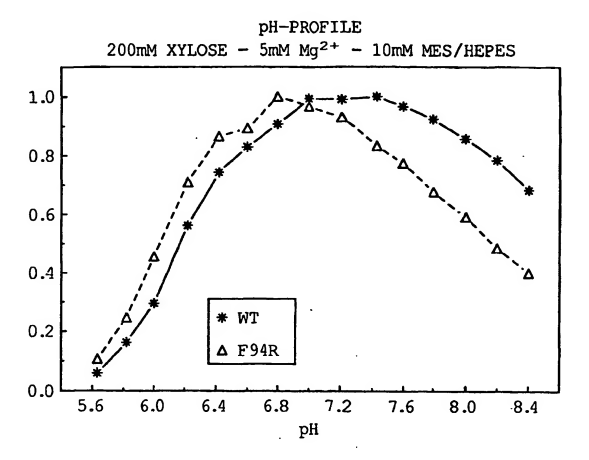


FIGURE 11

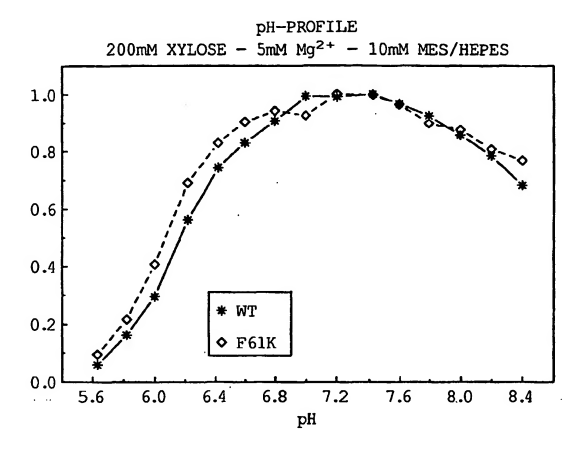


FIGURE 12

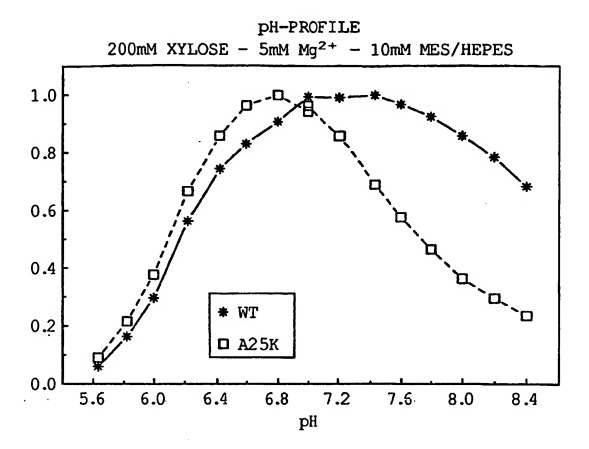


FIGURE 13

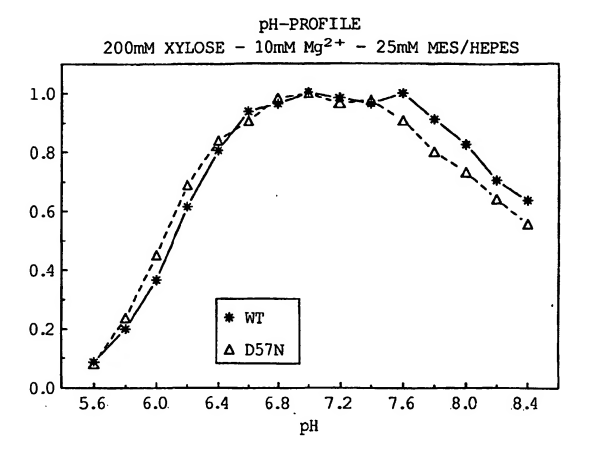


FIGURE 14

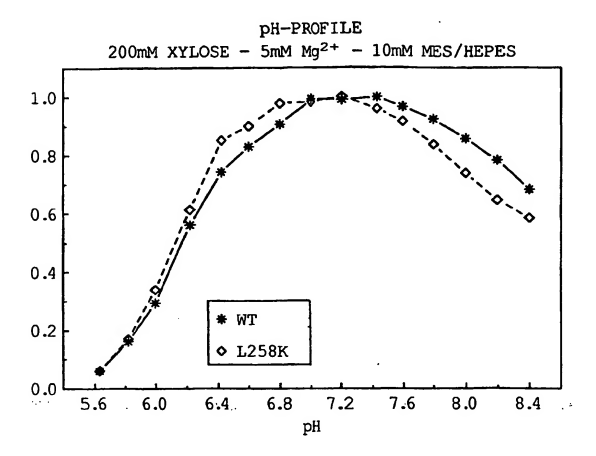


FIGURE 15

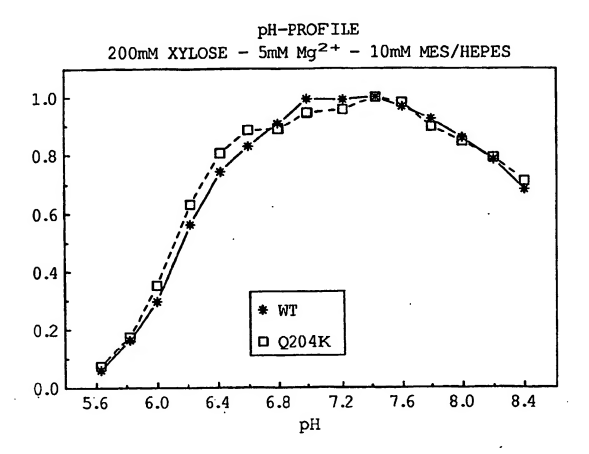


FIGURE 16

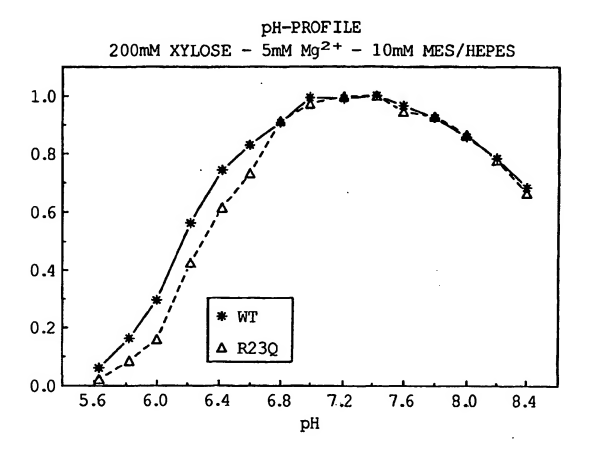


FIGURE 17-

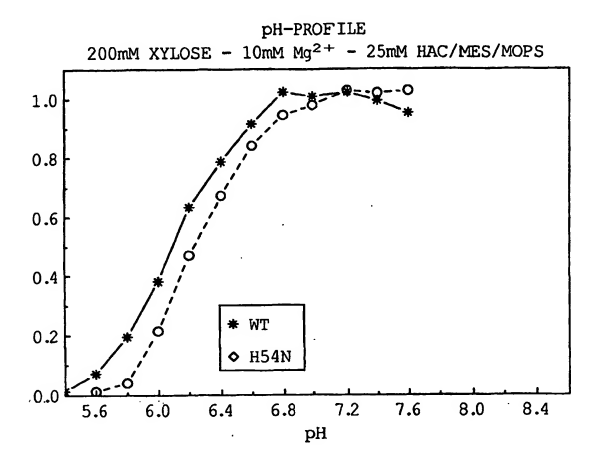
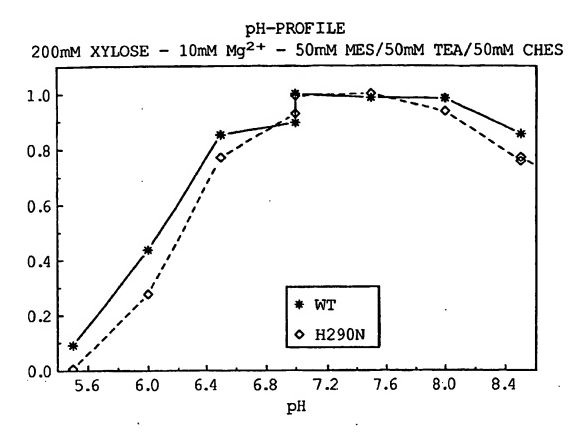


FIGURE \_18



.. FIGURE 19

pH-PROFILE 200mM XYLOSE - 5mM Mg<sup>2+</sup> 1.0 0.8 0.6 0.4 ◆ T95D 0.2 0.0 6.0 6.4 7.2 7.6 5.6 6.8 8.0 8.4 pН

FIGURE 20

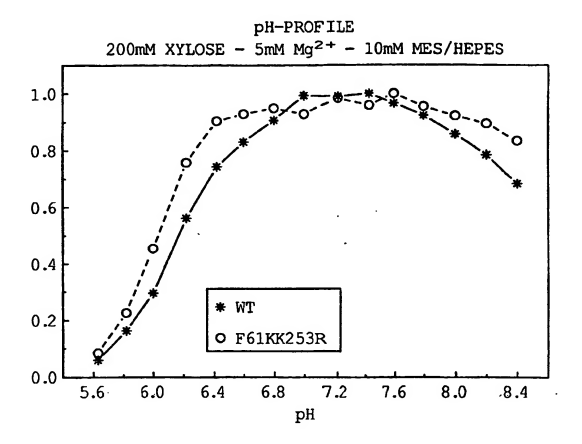


FIGURE 21